Emergency Medicine and Trauma Care Journal

Research Article

Todorova DG and Andonova AN. Emerg Med Trauma. EMTCJ-100029

Albumin or Hydroxyethyl Starch vs Saline for Volume Replacement in Surgical Patients partly Suggest Starling’s Law is Wrong, What Replacement is there? Perspective

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Citation: Ghanem AN (2020) Albumin or Hydroxyethyl Starch vs Saline for Volume Replacement in Surgical Patients partly Suggest Starling’s Law is Wrong, What Replacement is there? Perspective. Emerg Med Trauma. EMTCJ-100030

Received date: 19 March, 2020; Accepted date: 21 March, 2020; Published date: 27 March, 2020

Abstract

Introduction and objective: A recent article in JAMA demonstrated that the use of HES for volume replacement therapy, compared with 0.9% saline, resulted in no significant difference. This affirms that oncotic pressure of albumin is a fallacy in vivo, and Starling’s law is partly wrong. This perspective reports the complete evidence that Starling’s law is wrong, and the correct replacement is the hydrodynamic of G tube. The clinical relevance and significance is discussed.

Material and methods: The physics proof is based on G tube hydrodynamic. Physiological proof is based on study of the hind limb of sheep: running plasma and later saline through the artery compared to that through the vein as regards the formation of oedema. The clinical significance is based on 2 studies one prospective and a 23 case series on volumetric overload shocks (VOS).

Results: Hydrodynamics of G tube showed that proximal, akin to arterial, pressure induces suction “absorption” not “filtration”. In Poiseuille’s tube side pressure is all positive causing filtration based on which Starling proposed his hypothesis. The physiological evidence proves that the capillary works as G tube not Poiseuille’s tube: Oedema occurred when fluids are run through the vein but not through the artery. There was no difference using saline or plasma proteins in physiological and clinical studies. The wrong Starling’s law dictates the faulty rules on fluid therapy inducing VOS and ARDS.

Conclusion: Hydrodynamic of the G tube challenges the role attributed to arterial pressure as filtration force in Starling’s law. A literature review shows that oncotic pressure does not work either. The new hydrodynamic of G tube is proposed to replace Starling’s law which is wrong on both forces. The physiological proof and relevance to clinical importance on the pathogenesis of clinical syndromes are discussed.

Key points

Question: The use of HES for volume replacement therapy, compared with 0.9% saline, resulted in no significant difference. This means that neither HES nor albumin substitute has oncotic effect. What is the proof Starling’s law wrong and is there replacement and what is its clinical significance?

Findings: There is physics and physiological and clinical evidence to prove Starling’s law is wrong. There is evidence to suggest that using plasma or substitute of HES is no different from saline. This demonstrates oncotic pressure does not exist. Physics research demonstrates hydrostatic capillary (dynamic lumen) pressure induces suction not filtration. Physiological evidence demonstrates that the capillary works as G tube not Poiseuille’s tube. Hence...
Starling’s law is wrong on both hydrostatic and oncotic pressure forces. The correct replacement is the hydrodynamics of G tube.

**Meaning:** Starling’s law is wrong on both forces. The correct replacement is the hydrodynamics of the G tube. The clinical significance is discussed.

**Keywords:** AKI; ARDS; Albumin vs Saline; Capillary-interstitial fluid transfer; Capillary wall; Fluid therapy; Glycocalyx, Hyponatraemia; MODS; Pre-capillary sphincter; The TURP syndrome; Starling’s law; Shock; Volume kinetics

**Abbreviations**

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>ARDS</td>
<td>The adult respiratory distress syndrome</td>
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<td>AKI</td>
<td>Acute kidney injury</td>
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<td>BFT</td>
<td>Bolus Fluid Therapy</td>
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<td>CP</td>
<td>Chamber Pressure</td>
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<tr>
<td>CVS</td>
<td>Cardiovascular system</td>
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<td>CNS</td>
<td>Central Nervous system</td>
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<td>DP</td>
<td>Distal Pressure</td>
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<td>EGDT</td>
<td>Early Goal-Directed Therapy</td>
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<td>FP</td>
<td>Flow Pressure</td>
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<tr>
<td>G tube</td>
<td>Porous orifice tube</td>
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<tr>
<td>HN</td>
<td>Hyponatraemia</td>
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<tr>
<td>HST</td>
<td>Hypertonic sodium therapy of 5% NaCl and/or 8.4% Sodium Bicarbonate</td>
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<tr>
<td>ISF</td>
<td>Interstitial fluid</td>
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<tr>
<td>ICU</td>
<td>Intensive care unit</td>
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<tr>
<td>LP</td>
<td>Lumen Pressure</td>
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<tr>
<td>MODS</td>
<td>Multiple organ dysfunction syndrome</td>
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<tr>
<td>NaCo₃</td>
<td>Sodium bicarbonate</td>
</tr>
<tr>
<td>NaCl</td>
<td>Sodium chloride</td>
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<tr>
<td>PV</td>
<td>Plasma volume</td>
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<tr>
<td>PP</td>
<td>Proximal Pressure</td>
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<tr>
<td>RCT</td>
<td>Randomized controlled trial</td>
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<tr>
<td>TURP</td>
<td>The transurethral resection of the prostate</td>
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<tr>
<td>VK</td>
<td>Volume Kinetic</td>
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<tr>
<td>VO</td>
<td>Volumetric Overload</td>
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<tr>
<td>VOS</td>
<td>Volumetric overload shocks</td>
</tr>
<tr>
<td>VOS1</td>
<td>Volumetric overload shock, Type 1</td>
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<tr>
<td>VOS2</td>
<td>Volumetric overload shock, Type 2</td>
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<td>WW2</td>
<td>World War Two</td>
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**Introduction**

The authors of this article by [1] on the Effect of Hydroxyethyl Starch vs Saline for Volume Replacement Therapy on Death or Postoperative Complications Among High-Risk Patients Undergoing Major Abdominal Surgery: The FLASH Randomized Clinical Trial in *JAMA* are commended on their excellent powerful study and report. It is a true representation of evidence based medicine. It has inspired me to write this perspective. This study would be perfect had they included one piece of data in the result section: The volumetric balance of patients during the surgery time i.e. volumetric overload over time (VO/T) that is compared for the occurrence of morbidity and mortality of the two groups receiving Hydroxyethyl starch (HES) vs saline. There should be a significant difference between patients with morbidity and mortality, and those who do not have such complications.

I believe the authors can retrieve these data and subject it to statistical analysis if they would. This will make their article the first ever prospective study to report and incriminate volumetric overload (VO) of the given HES and saline in surgical patients who suffer morbidity of acute respiratory distress syndrome (ARDS), multiple organ dysfunction syndrome (MODS), and/or acute kidney injury (AKI) and mortality after major surgery. The patients of the two groups receiving HES or saline should be divided into 3 subgroups each of asymptomatic (A), symptomatic with morbidities (B) and the dead or mortality (C). The mean volume of retained fluid VO is calculated for each of the subgroups and statistically compared. I dare say that I anticipate a high statistical significance (p=0.0001) as my research demonstrated (Figure 1,2 Table 1).

**Figure 1:** Shows the means and standard deviations of volumetric overload in 10 symptomatic patients presenting with shock and hyponatraemia among 100 consecutive patients during a prospective study on the transurethral resection of the prostate (TURP). The fluids were of Glycine absorbed (Gly abs), intravenously infused 5% Dextrose (IVI Dext) Total IVI fluids, Total Sodium-free fluid gained (Na Free Gain) and total fluid gain in litres. (Reproduced with permission of the author and Editor of BJU Int. from this article reference 8).
Figure 2: Shows volumetric overload (VO) quantity (in litres and as per cent % of body weight) and types of fluids. Group 1 was the 3 patients who died in the case series as they were misdiagnosed as one of the previously known shocks and treated with further volume expansion. Group 2 were 10 patients from the series who were correctly diagnosed as volumetric overload shock and treated with hypertonic sodium therapy (HST). Group 3 were 10 patients who were seen in the prospective study and subdivided into 2 groups; Group 3.1 of 5 patients treated with HST and Group 3.2 of 5 patients who were treated with guarded volume expansion using isotonic saline. (Reproduced with permission of the author from this article reference 21).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Std. Err</th>
<th>Std. Value</th>
<th>T Value</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>-</td>
<td>-</td>
<td>-0.773</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Fluid Gain (l)</td>
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<td>0.228</td>
<td>1.044</td>
<td>3.721</td>
<td>0.0001</td>
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<td>Osmolality</td>
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<td>-0.375</td>
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<td>0.0212</td>
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<td>Na+ (C_B)</td>
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<td>0.049</td>
<td>0.616</td>
<td>1.95</td>
<td>0.0597</td>
</tr>
<tr>
<td>Alb (C_B)</td>
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<td>0.087</td>
<td>0.239</td>
<td>0.713</td>
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<tr>
<td>Hb (C_B)</td>
<td>-0.282</td>
<td>0.246</td>
<td>-0.368</td>
<td>1.149</td>
<td>0.2587</td>
</tr>
<tr>
<td>Glycine (C_B)</td>
<td>-4.97E-05</td>
<td>5.98E-05</td>
<td>-0.242</td>
<td>0.832</td>
<td>0.4112</td>
</tr>
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</table>

Table 1: Shows the multiple regression analysis of total per-operative fluid gain, drop in measured serum osmolality (OsmM), sodium, albumin, Hb and increase in serum glycine occurring immediately post-operatively in relation to signs of the TURP syndrome. Volumetric gain and hypo-osmolality are the only significant factors; fluid gain is highly significant. (Reproduced with permission of the author and Editor of BJU Int. from this article reference 8).

The conclusion of the study is: Among patients at risk of postoperative kidney injury undergoing major abdominal surgery, use of HES for volume replacement therapy, compared with 0.9% saline, resulted in no significant difference in a composite outcome of death or major postoperative complications within 14 days after surgery.” This means that HES like albumin has no oncotic effect in vivo as has previously been proved [2]. This is not surprising at all as the HES fluids were introduced as plasma substitutes. Furthermore, it affirmed with powerful evidence based medicine article that half the equation of Starling’s law on the capillary-interstitial fluid transfer concerning the oncotic pressure of plasma albumin causing absorption is wrong. My research demonstrated that the other half of this law on hydrostatic pressure causing filtration across the capillary wall is also wrong. The correct replacement for this law is hydrodynamic of the porous orifice (G) tube as explained here.

Professor Fifner’s article at BMJ on “saline versus albumin fluid evaluation (SAFE) 20062, concluding: “saline or albumin produces similar outcome”. In 1998 meta-analysis, albumin faired worse3, justifying BMJ slogan “Why albumin may not work”. This is conflicting and perplexing. Professor Vincent [3] mentioned in his editorial at BMJ: “the aim of the analysis was to show that albumin administration is safe.” I wish all the luck with the “Save Albumin Campaign”, and confirm that I do not deny albumin safety and usefulness when correctly indicated. The valid painstaking analysis of data and the conclusion of similar or worse outcome, to my mind, re-affirms the fact that albumin oncotic pressure in VIVO is fallacy [4-6], explaining BMJ slogan. It is shameful waste to spend so much effort and money on huge clinical trials, on the wrong basic notion that albumin oncotic pressure exists in VIVO. My aim here, however, is to discuss issues overlooked in all SAFE [2] and FLASH [1] trials data analysis highlighting a
Volumetric overload (VO) over Time (VO/T) is a concept verifiable by comparing patients' body weight on ICU to that on hospital admission or before and immediately after surgery. This reveals a staggering VO of 12-14 litres of retained fluid in the acute respiratory distress syndrome (ARDS) patients! In 1967, Professor [7] documented this volumetric fluid gain in the first report on ARDS, which became later known as the multiple organ dysfunction syndrome (MODS). Such VO data have not ever been since documented in any report other than mine. My research demonstrated that multiple regression analysis has proved that volumetric overload is the most significant factor in causing the clinical picture of VOS1 of the TURP syndrome [8] (Figure 2, 3 and Table 1). Volumetric balance is consistently missed in prospective trials. Not a single prospective SAFE or other trial report volumetric data in MODS syndrome patients, ARDS or AKI!

Fluid Type and Volume, and Time of gain, have vital significance in the pathogenesis and outcome of MODS or ARDS in post-surgical patients. Type of fluid gives characteristic serum solute dilution markers. Volume is directly, while time is inversely, related to the severity. Sodium-free fluids (Type 1) or VO1 dilute all serum or extracellular fluid contents including albumin, but its best marker is hyponatraemia (HN). The well-known transurethral resection of the prostate (TURP) [8] syndrome is a “model” of many such cases of HN of albumin or crystalloids seen in clinical practice. This “model” of TURP syndrome means it can be, and has been, precisely reproduced in animals in the absence of sepsis, hypothermia and recognized shocks. Hyponatraemia of <120 mmol/l is common hospital iatrogenic complication of fluid therapy that affects men, women and children and is usually lethal.

The TURP syndrome is induced by both the irrigating fluid absorption (1.5% Glycine, Sorbitol or Mannitol) and the infused intravenous fluids such as 5% Glucose [9, 10]. A quantity of 3.5-5 litres (l), gained during 1 hour surgery, induces a classical condition of volumetric overload shock type 1 (VOS1) while 5-6l may be lethal.8 The VO of 3.5l may be considered normal daily intake and is tolerated over couple of hours but when gained in one hour it becomes pathological. The condition manifests clinically with paradoxical hypotension shock (Paradoxical means hyper-NOT hypo-volaemic shock) with features unrecognizable from or identical to hypovolaemic shock except for bradycardia and transient rare elevation of arterial pressure. It also has paradoxical AKI among other features of the MODS syndrome. This must to be kept in mind in order to recognize VO/T of VOS2, induced by SAFE fluids, with scarce markers if any. This is important as the TURP procedure is currently performed in saline irrigation (TURIS), so much more VOS2 with scarce or no markers will soon appear, and is also common in patients undergoing major surgery who received overzealous saline, HES or plasma infusions.

The common thought and practice of treating physician in such paradoxical VO/T shock is to infuse further volume of either SAFE albumin or crystalloids isotonic fluid! The physician aims
to elevate arterial pressure by increasing vascular volume in the belief that he/she is facing hypovolaemic hypotension shock, while data indicate VO/T shock. The action just makes it worse or irreversible shock and establishes MODS when the patient is shifted to ICU. The insult of both SAFE and crystalloids isotonic fluids may occur in resuscitating the TURP syndrome with definite characteristic serum markers and proven clinical features, or may complicate overzealous resuscitation of any recognized shock, trauma, during major surgery or in ICU patients when serum markers are scarce or nil. Nothing to guide physician at all except his thought determined by current basic teaching on vascular volume and pressure relationship on one hand, and the forces of Starling’s law regulating the capillary-ISF circulation on the other. The latter determine the type and volume of SAFE fluid used in resuscitation of shock, trauma, burns, haemorrhage and sepsis and prolonged major surgery. So the faulty Starling’s law is responsible for misleading physicians into giving too much fluids.

Sodium-based fluids (VO2) such as saline, albumin, plasma substitutes of HES and blood may also induce VO/T shock identified as volumetric overload shock type 2 (VOS2). It may complicate the resuscitation of the TURP syndrome when VO2fluids erase HN while worsening VO. The main serum marker of using saline or HES becomes hypoalbuminaemia [9] that is not as marked as HN. It also has the same clinical features of paradoxical hypotension shock and MODS Syndrome. It may complicate resuscitation of any recognized shock and fluid resuscitation during major surgery complicating both saline and plasma or substitutes of HES. The transition from hypo- to hyper-volaemic hypotension shock is hard or impossible to detect. No stop sign to show that such patient is having hyper-volaemic not hypo-volaemic hypotension shock. None to warn when the quantity needed in treating true hypovolaemia is surpassed. Vascular pressures of central venous pressure (CVP), pulmonary capillary wedge pressure (PCWP) and blood pressure (BP) changes of VO/T are identical except for bradycardia and an occasional transient initial rise of BP10. Massive plasma and blood infusions have no serum markers at al or specific vital signs except VO2 increase of body weight or calculated VO and occurrence of MODS or ARDS or AKI occurring in any combinations, but one system may predominate.

It should be realized that hypotension is not always synonymous with hypovolaemia. It is worth mentioning also that, up to this point, sepsis is as innocent as the wolf in Joseph story. A little later, sepsis will do its nasty work and further complicate MODS into its current trendy name associated with sepsis, termed the systemic inflammatory response syndrome (SIRS). The scientific basis that underlies physician’s thought while resuscitating a patient is explained later. The evidence on how and why Starling’s law10 is wrong on both forces while continuing to dictate the faulty rules on fluid therapy concerning the type and incorrect volume of fluid in resuscitation while volume is consistently missed in SAFE trials is explained here.

On the physiological issues; the direct positive relationship of fluid volume and pressure, the work of Poiseuille on flow and pressure exerted on the wall of strait uniform brass tubes, as well as the albumin oncotic pressure, were all imported by Starling10 in 1896 direct from physics to medicine at the Lancet without any physiological verification or testing what so ever! Modern clinical chemistry allowed verification of albumin oncotic pressure by [6] comparing variety of body fluids to plasma protein- there was no difference. The real ultra-structure of the capillary wall revealed by Karnovsky [5] and the pre-capillary sphincter revealed by Rhodin [11] were reported in 1967. Hence Renkin called for reconsideration of Starling’s hypothesis [12] in 1986. The hydrodynamic of a porous orifice tube was reported in 2001[13]. Thus, all physiological research done before 1962 that advocated Starling’s hypothesis promoting it into law is invalid. Based on the consequences of capillary permeability to macromolecules demonstrating that oncotic pressure does not work in vivo, Renkin [12] advocated reconsideration of Starling’s hypothesis. What alternative was there then? There was none. Only an idea in mind derived from clinical observation on the use of fluids in resuscitation of shock, trauma and the TURP syndrome was communicated and reported at BMJ9 in 1985. The results of the physics study on the hydrodynamic of the G tube were completed but unreported until 2001 [13].

The direct proportional relationship of fluid volume to pressure works in the vascular system up to a limit only. This is true in physics too, if too much fluid is pushed into a reservoir above its capacity, it will burst and the volume-pressure relationship vanishes. Thus, perhaps volume replacement in shock should not exceed in total the maximum capacitance of vascular system of 7l in adult. Considering that blood loss is fatal when about half the vascular volume is acutely lost, a replacement should not exceed the lost volume after control of bleeding. After any overzealous vascular volume expansion, the excess fluid must leak out into and drown the interstitial space and cells! The most deleterious effect of such internal drowning is on the vital organs. Both vital organ signs8 and post-mortem findings [14] demonstrate the massive volume of retained fluids. This letter to Editor [14] is the only documented evidence in literature that reported the massive retained fluid volume with swollen vital organs at post mortem examination! The only article that reported retained fluid volume was the first report on ARDS by [7] in addition to the articles of mine some of which are referenced here.

Albumen oncotic pressure, no doubt, exists in vitro across membrane impermeable to its molecules. Even, in such physics experiments, oncotic pressure is too weak and too slow force to be effectively and solely responsible for fluid return into capillary lumen6. It has cell building nutritional value, how does albumin reach the cells? It must cross the capillary wall! However, the evidence that oncotic pressure works in vivo is non-existing [1-3]. The only difference between albumen or HES and saline fluids in SAFE and FLASH trials is the added albumen or starch presumed
to have oncotic pressure, a function of its molecule size in relation to pore size or permeability of membrane. The pores of normal capillary wall became known [8] decades after Starling’s report on his hypothesis. Karnovesky has shown the intercellular slits between capillary wall cells to allow horse radish, a much larger molecule than albumin, to pass freely across the capillary wall [5]. As the result of these trials 1,2 demonstrated that both SAFE fluids have similar outcome, this further re-affirms that albumen oncotic pressure in clinical medicine [1,2-4], clinical chemistry [6] and modern physiology [15-17] is fallacy in vivo, simply because albumen molecules pass freely across the large pores of normal capillary membrane [5].

This may answer the BMJ slogan: Why albumin may not work. So, irrespective whether albumin has equal or worse outcome, the fact that it did not show clear superiority to saline in SAFE trials is affirmative evidence that albumen oncotic pressure is fallacy in vivo. Such fallacy has also been long proved in biochemical research [5] and physiological research [15-17]. Oncotic pressure is the presumed main absorption force in capillary-interstitial fluid (ISF) transfer, and represents one half of the equation of Starling’s law [10]. Thus as it has proved wrong, the law must be wrong! This was the reason for the call to reconsider Starling’s hypothesis [12]. However, there was no existing alternative then- only an idea in mind communicated at BMJ in 1985. This was later verified and reported, the clinical work [8] in 1990 and physics work [13] in 2001 as well as in more recent articles [18-21], and a book [22].

Verifying the other half of Starling law equation concerning capillary arterial pressure as the filtration force was my objective. Does the capillary have positive pressure on its wall pushing fluid out? Does the flow pressure akin to arterial pressure cause filtration? Does the capillary tube act like Poiseuille’s strait uniform tube and have positive pressure on its wall that pushes fluid out through pores? The ultra-structure of capillary wall [5] and pre-capillary sphincter [11] were discovered 80 years after Starling reported his hypothesis. The capillary proved a porous orifice tube. I made several porous tubes fitted with narrow orifice mimicking the capillary with a pre-capillary sphincter on a larger scale to verify this. The G tube was surrounded by a chamber C to represent the ISF space. These porous orifice tubes were used to study hydrodynamic flow and pressure and compared to Poiseuille’s tube, particularly in relation to the side pressure exerted on its wall.

The porous orifice (G) tube hydrodynamic proves totally different to Poiseuille’s tube. There was no positive pressure exerted on the wall and no fluid filtered out over the proximal half of the porous orifice tube. The flow pressure representing arterial pressure is not responsible for filtration! It caused mainly suction maximum at the proximal part of the G tube. Thus the main force in the equation on Starling’s law concerning arterial capillary pressure filtration is also wrong. How does it work? What pushes the fluid out and what returns it in? Does the G tube hydrodynamic offer a complete hypothesis to explain the capillary-interstitial fluid exchange? How does it relate to physiology and medicine? To know the answer to these questions on a most fascinating phenomenon of the G tube, please read the articles [12,17-21] or the book [22].

So, the law dictating the scientific basis that underlies physician’s thought on vascular volume expansion at resuscitation of shock, trauma and acutely ill patients is wrong on all accounts. The most harmful part of this erroneous law is in fact that concerning arterial pressure, presumed to be the main filtration force in the capillary. This is the part that Starling thought the capillary acts like Poiseuille’s strait uniform tube, exerting positive pressure on the wall that filters fluid out.

The complete evidence that Starling’s law is wrong and the correct replacement is the hydrodynamic of the porous orifice tube is summarised here, and has been reported elsewhere [23]:

Dr Starling proposed his hypothesis >80 years prior to the discovery of the capillary ultrastructure and correct physiology which are as follows. He based his hypothesis on Poiseuille’s work [24] in which the hydrostatic pressure is a positive function of the arterial pressure causing filtration, but in the G tube hydrodynamic as a porous orifice tube- akin to the capillary this pressure is different causing suction. Thus Starling’s law is wrong on both forces because:

- The capillary has a pre-capillary sphincter as reported by Rhodin in 1967 [11] which makes it different from Poiseuille’s tube of uniform diameter as my research demonstrated.
- The capillary has porous wall of intercellular slits that allow the passage of plasma proteins as shown by Karnoveski in 1967. Hence plasma proteins cannot exert an oncotic pressure in vivo.
- The osmotic chemical composition of various body fluids is identical to plasma proteins as demonstrated by Hendry in 19626. Hence oncotic pressure if it exists is too week and too slow to cause absorption.
- The oncotic pressure of plasma proteins does not work as absorption force neither in physiology as proved by Hendry in 19626 nor in clinical practice demonstrated by Cochrane Injuries Group in 1998.
- More recent evidence demonstrates that both plasma proteins and HES1 versus saline show no significant difference during fluid infusion for resuscitation of acutely ill patients and those undergoing major surgery.
- Guyton and Coleman (1968) demonstrated that ISF space has a negative pressure [25] of -7 cm water and [26] showed that the lymph has the same negative pressure. The pressure under the skin is negative. That cannot be explained by Starling's forces.
- Inadequacy in explaining the capillary--ISF transfer in many parts of the body as reported by [27,28], particularly vital organs, has previously called for reconsideration of Starling’s hypothesis by Renkin in 1986 [12].

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Emerg Med Trauma, an open access journal
ISSN: 2652-4422
My physics and physiological research work has demonstrated that the hydrostatic or rather lumen dynamic “arterial” pressure does not cause filtration across the wall of porous orifice (G) tube as proposed by Starling. It causes suction.

This lumen pressure (LP) induces negative side pressure gradient along the G tube causing suction maximum near the inlet and turns positive maximum near the exit causing filtration as based on physics experiments and physiological research. Venous pressure enhances filtration and causes oedema but arterial pressure does not- it causes absorption by suction.

The physiological study on the hind limb of sheep has completed the evidence that Starling’s law is wrong as the capillary works as G tube not Poiseuille’s tube.

Starling’s law being wrong underlies all errors and misconceptions on fluid therapy misleading physicians into giving too much fluid during resuscitation of shock and the acutely ill patients and during prolonged surgery inducing VO/T and causing ARDS.

Received thinking that elevating central venous pressure (CVP) is synonymous with elevating arterial pressure is prevailing in current clinical practice during fluid therapy for shock and the management of the acutely ill patients, but wrong. This is undoubtedly correct during restoration therapy for hypovolemic and haemorrhagic shock, but vascular expansion or volumetric overload (VO) is a different issue as it induces VOS and ARDS.

Persistent attempts to elevate CVP up to levels of 18 to 22 cm water are common received practice. The normal CVP, however, is around 0 and most textbooks report a range of -7 to +7 cm water.

Clinical observations demonstrate that, in addition to the well-known effect of high venous pressure causing oedema, arterial hypertension has no such effect, if not the exact opposite. In clinical practice, although arterial hypertension is common, ISF oedema is unknown among its complications.

In the G-C model, a minor increase in DP increases fluid volume in chamber C around the G tube reverting CP from negative to positive while slowing the G-C circulation. Increasing DP has similar effect to decreasing PP on the G-C circulation and chamber pressure and volume.

Vascular expansion causes VO shocks. There is no doubt that the erroneous Starling’s law is responsible for the many errors and misconceptions prevailing on fluid therapy for shock and the acutely ill patients which mislead physicians into giving too much fluid that induce volumetric overload shocks (VOS) causing MODS or ARDS.

This underlies the treating physician’s thought when embarking on overzealous fluid infusion during the resuscitation of shock and prolonged major surgery. He was taught that volume expansion has direct positive unlimited relationship with pressure. It is the only way he knows off to improve capillary circulation. Well, it does not. Volume replacement is effective when an actual blood volume loss is restored to normal that is less than maximum capacity of the vascular system. After that the relation of volume to pressure is reversed. Any excess volume, vascular expansion or hypervolaemia of VO/T induces hypotension shock just like hypovolaemia does! Considering the concept of VO/T by reporting volume of fluids in future SAFE trials and verifying the scientific basis of fluid resuscitation in shock are needed for resolving the puzzle of MOVD/F syndrome or ARDS and improving outcome of surgical patients on ICU.

References


